PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11	(11) International Publication Number: WO		
A61K 7/02, 7/48	Al	(43	3) International Publication Date:	29 October 1998 (29.10.98)	
(21) International Application Number: PCT/IBs (22) International Filing Date: 30 March 1998 (3)		- 1	(81) Designated States: AU, CA, CN, patent (AT, BE, CH, DE, DK, LU, MC, NL, PT, SE).	CZ. IP, MX. US. European ES, FI. FR. GB, GR, IE, IT,	
(30) Priority Data: 9708051.9 22 April 1997 (22.04.97)	G	В	Published With international search repor	rı.	
(71) Applicant (for all designated States except US): THE TER & GAMBLE COMPANY [US/US]; One Pi Gamble Plaza, Cincinnati, OH 45202 (US).	PROC rocter	C.			
(72) Inventors; and (75) Inventors/Applicants (for US only): BELL, Stephe [IE/GB]; 65 Ferney Road, E. Barnett, Herts EN4 8L BOXSHALL, Alison, Ruth [GB/GB]; 153A Churc Staines, Middlesex TW18 4XZ (GB).	LB (GB). l			
(74) Agents: REED, T., David et al.; The Procter & Company, 5299 Spring Grove Avenue, Cincinn 45217 (US).	Gambl ati, O	le H			
(54) Title: COSMETIC COMPOSITIONS					

(57) Abstract

Cosmetic composition comprising: (a) from about 0.1 % to about 20 % by weight of particulate nylon; and (b) from about 0.1 % to about 20 % by weight of a liquid, polyol carboxylic acid ester having a polyol moiety and at least 4 carboxylic acid moities, wherein the polyol moiety is selected from sugars and sugar alcohols containing from about 4 to about 8 hydroxyl groups, and wherein each carboxylic acid moiety has from about 8 to about 22 carbon atoms, and wherein said liquid polyol carboxylic acid ester has a complete melting point of less than about 30 °C.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	. Slovenia
AM	Armenia	Ħ	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑÜ	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
۸Z	Azerbaijan	GB	United Kingdom	мс	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The fermer Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Muli	11	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	ÜA	Ukraine
BR	Brazil	IL.	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	17	Italy	MX	Mexico	UZ	Uzhekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Vict Nam
CG	Congo	KE	Kenya	NL.	Netherlands	YU	Yogoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
a	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand	2**	STUDENCE
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Pederation		
DE	Germany	ш	Liechtenstein	50	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG			
			La OCT (4	3G	Singapore		

Field of the Invention

The present invention relates to cosmetic compositions and more particularly, to pigmented foundation make-up compositions and concealers.

Background of the Invention

A foundation composition can be applied to the face and other parts of the body to even skin tone and texture and to hide pores and imperfections, and the like. A foundation composition is also applied to moisturize the skin and to balance the oil level of the skin. Make-up compositions are generally available in the form of liquid or cream suspensions, emulsions, gels, pressed powders or anhydrous oil and wax compositions. Such cosmetic make-up compositions are described in US-A-3,444,291, US-A-4,486,405, US-A-4,804,532, US-A-3,978,207, US-A-4,659,562, US-A-5,143,722 and Nakamura et al., Preprints of the XIVth I.F.S.C.C. Congress, Barcelona, 1986, Vol. I, 51-63 (1986).

Skin is subject to abuse by many extrinsic (environmental) factors as well as intrinsic (chronoaging) factors. A common extrinsic factor is exposure to ultraviolet radiation. Whether extrinsic or intrinsic, the abuse results in wrinkling of the skin. To many people, skin wrinkles are a reminder of the disappearance of youth. Attempts have therefore been made to reduce the appearance of wrinkles. These treatments range from cosmetic creams and moisturizers to various forms of cosmetic surgery. However, these treatments have not always been successful. In particular, attempts to cover up wrinkles using cosmetic foundations often result in an undesirable accentuation of the appearance of wrinkles. In addition, for many people it is undesirable to resort to cosmetic surgery in order to reduce the appearance of wrinkles. It would therefore be desirable to provide a cosmetic composition, and in particular a foundation

1

COSMETIC COMPOSITIONS

composition which significantly reduces the appearance of wrinkles and fine lines.

Particulate nylon is a polyamide polymer and is known in the art for use in cosmetic compositions. In particular, nylon particles are known for use in cosmetic emulsion compositions as described in "Active Delivery from Nylon Particles", Cosmetic and Toiletries, Dec 1993, Vol. 108, pages 97-100, and "Nylon Powders in Cosmetics", Manufacturing Chemist, May 1984, pages 73-75.

EP-A-0592107 (published 13th April 1994) discloses a powdered cosmetic composition comprising ultrafine powdered boron nitride and optionally powdered nylon. EP-A-0462709 (published 27 December 1991) discloses a cosmetic composition comprising light oils, cosmetic waxes and a powder phase comprising spherical silica, particulate nylon and spherical powder.

US95/15375 and US95/15374 (The Procter & Gamble Company) relates to cosmetic compositions comprising liquid polyol carboxylic acid esters and which in addition can contain particulate materials such as silica, titanium dioxide, zinc oxide and iron oxide.

It has now surprisingly been found that by incorporating particulate nylon into a cosmetic composition together with a liquid, polyol carboxylic acid ester a cosmetic composition is provided which exhibits significant reduction in the appearance of fine lines and wrinkles.

It is accordingly an object of the present invention to provide a cosmetic composition having improved reduction in the appearance of fine lines and wrinkles when applied to the skin.

It is a further object of the present invention to provide a cosmetic composition which delivers excellent coverage, application and skin feel characteristics.

3

Summary of the Invention

In accordance with one aspect of the present invention, there is provided a cosmetic composition comprising:

- (a) from about 0.1% to about 20% by weight of particulate nylon; and
- (b) from about 0.1% to about 20% by weight of a liquid, polyol carboxylic acid ester having a polyol moiety and at least 4 carboxylic acid moieties, wherein the polyol moiety is selected from sugars and sugar alcohols containing from about 4 to about 8 hydroxyl groups, and wherein each carboxylic acid moiety has from about 8 to about 22 carbon atoms, and wherein said liquid polyol carboxylic acid ester has a complete melting point of less than about 30°C.

The cosmetic compositions of the present invention provide a significant reduction in the appearance of fine lines and wrinkles when applied to the skin.

According to a second aspect of the present invention there is provided the use of such compositions for reducing the appearance of fine lines and wrinkles when applied to the skin.

All levels and ratios are by weight of total composition, unless otherwise indicated. Chain lengths and degrees of alkoxylation are also specified on a weight average basis.

Detailed Description of the Invention

The cosmetic composition according to the present invention comprises particulate nylon and a liquid, polyol carboxylic acid ester.

Nylon

A first essential component of the cosmetic compositions of the invention is particulate nylon. The particulate nylon is present in the compositions of the invention at a level of from about 0.1% to about 20%, preferably

WO 98/47470

from about 0.1% to about 10%, more preferably from about 4% to about 8% by weight of composition.

4

Nylon is a water-insoluble, polyamide formed from the reaction of a diamine and a di-acid halide. Several different types of nylon can be formed by altering the carbon chain length or configuration in the amine or diacid. Nylon can be represented by the general formula HN-(CH₂)_n-C_n=O. In general n lies in the range of from about 6 to about 12. Examples of nylon compounds encompassed by this general formula are nylon-6 and nylon-12.

Suitable particulate nylon materials for use in the compositions of the present invention include those materials having a spheroidal, porous structure. Without wishing to be bound by theory it believed that it is the particulate, spheroidal, porous nature of the nylon materials which interacts with the liquid polyol carboxylic acid ester skin conditioning agent to provide a reduction in the appearance of fine lines and wrinkles. Hence according to a further aspect of the present invention there is provided a cosmetic composition comprising a spheroidal, porous particulate material and a liquid polyol carboxylic acid ester. Preferred for use herein are particulate, spheroidal, porous nylon materials having a volume average particle size of from about 5 microns to about 30 microns, preferably from about 10 microns to about 20 microns.

Particularly preferred from the viewpoint of reducing the appearance of fine lines and wrinkles is nylon 12. This is commercially available from Chesham Chemicals, Cunningham House, Westfield Lane, Harrow, Middlesex, HA3 9ED under the tradename Orgasol 2002 series. A particularly preferred nylon for use herein is Orgasol 2002 EX D NAT.

Polyol Ester Skin Conditioning Agent

A second essential ingredient for use herein is a polyol ester skin conditioning agent.

The compositions of the present invention preferably comprise from about 0.01% to about 20%, more preferably from about 0.1% to about 15%, and especially from about 1% to about 10% by weight of the polyol ester.

The polyol ester preferred for use herein is a nonocclusive liquid or liquifiable polyol carboxylic acid ester. These polyol esters are derived from a polyol radical or moiety and one or more carboxylic acid radicals or moieties. In other words, these esters contain a moiety derived from a polyol and one or more moieties derived from a carboxylic acid. These carboxylic acid esters can also be derived from a carboxylic acid. These carboxylic acid esters can also be described as liquid polyol fatty acid esters, because the terms carboxylic acid and fatty acid are often used interchangeably by those skilled in the art.

The preferred liquid polyol polyesters employed in this invention comprise certain polyols, especially sugars or sugar alcohols, esterified with at least four fatty acid groups. Accordingly, the polyol starting material must have at least four esterifiable hydroxyl groups. Examples of preferred polyols are sugars, including monosaccharaides and disaccharides, and sugar alcohols. Examples of monosaccharides containing four hydroxyl groups are xylose and arabinose and the sugar alcohol derived from xylose, which has five hydroxyl groups, i.e., xylitol. The monosaccharide, erythrose, is not suitable in the practice of this invention since it only contains three hydroxyl groups, but the sugar alcohol derived from erythrose, i.e., erythritol, contains four hydroxyl groups and accordingly can be used. Suitable five hydroxyl groupcontaining monosaccharides are galactose, fructose, and sorbose. Sugar alcohols containing six -OH groups derived from the hydrolysis products of sucrose, as well as glucose and sorbose, e.g., sorbitol, are also suitable. Examples of disaccharide polyols which can be used include maltose, lactose, and sucrose, all of which contain eight hydroxyl groups.

Preferred polyols for preparing the polyesters for use in the present invention are selected from the group consisting of erythritol, xylitol, sorbitol, glucose, and sucrose. Sucrose is especially preferred.

The polyol starting material having at least four hydroxyl groups is esterified on at least four of the -OH groups with a fatty acid containing from about 8 to about 22 carbon atoms. Examples of such fatty acids include caprylic, capric, lauric, myristic, myristoleic, palmitic, palmitoleic, stearic, oleic, ricinoleic, linoleic, linolenic, eleostearic, arachidic, arachidonic, behenic, and erucic acid. The fatty acids can be derived from naturally occurring or synthetic fatty acids; they can be saturated or unsaturated, including positional and geometrical isomers. However, in order to provide liquid polyesters preferred for use herein, at least about 50% by weight of the fatty acid incorporated into the polyester molecule should be unsaturated. Oleic and linoleic acids, and mixtures thereof, are especially preferred.

The polyol fatty acid polyesters useful in this invention should contain at least four fatty acid ester groups. It is not necessary that all of the hydroxyl groups of the polyol be esterified with fatty acid, but it is preferable that the polyester contain no more than two unesterified hydroxyl groups. Most preferably, substantially all of the hydroxyl groups of the polyol are esterified with fatty acid, i.e., the polyol moiety is substantially completely esterified. The fatty acids esterified to the polyol molecule can be the same or mixed, but as noted above, a substantial amount of the unsaturated acid ester groups must be present to provide liquidity.

To illustrate the above points, a sucrose fatty triester would not be suitable for use herein because it does not contain the required four fatty acid ester groups. A sucrose tetra-fatty acid ester would be suitable, but is not preferred because it has more than two unesterified hydroxyl groups. A sucrose hexa-fatty acid ester would be preferred because it has no more than two unesterified hydroxyl groups. Highly preferred compounds in which all the hydroxyl groups are esterified with fatty acids include the liquid sucrose octa-substituted fatty acid esters.

The following are non-limiting examples of specific polyol fatty acid polyesters containing at least four fatty acid ester groups suitable for use in the present invention: glucose tetraoleate, the glucose tetraesters of

soybean oil fatty acids (unsaturated), the mannose tetraesters of mixed soybean oil fatty acids, the galactose tetraesters of oleic acid, the arabinose tetraesters of linoleic acid, xylose tetralinoleate, galactose pentaoleate, sorbitol tetraoleate, the sorbitol hexaesters of unsaturated soybean oil fatty acids, xylitol pentaoleate, sucrose tetraoleate, sucrose pentaoletate, sucrose hexaoleate, sucrose hepatoleate, sucrose octaoleate, and mixtures thereof.

As noted above, highly preferred polyol fatty acid esters are those wherein the fatty acids contain from about 14 to about 18 carbon atoms.

The preferred liquid polyol polyesters preferred for use herein have complete melting points below about 30°C, preferably below about 27.5°C, more preferably below about 25°C. Complete melting points reported herein are measured by Differential Scanning Calorimetry (DSC).

The polyol fatty acid polyesters suitable for use herein can be prepared by a variety of methods well known to those skilled in the art. These methods include: transesterification of the polyol with methyl, ethyl or glycerol fatty acid esters using a variety of catalysts; acylation of the polyol with a fatty acid chloride; acylation of the polyol with a fatty acid anhydride; and acylation of the polyol with a fatty acid, per se. See U.S. Patent No. 2,831,854; U.S. Patent No. 4,005,196, to Jandacek, issued January 25, 1977; U.S. Patent No. 4,005,196, to Jandacek, issued January 25, 1977.

Optional ingredients

A wide variety of optional ingredients can be incorporated into the compositions. The following are non-limiting examples of numerous ingredients that can be used.

Oil Phase

The composition of the present invention can be formulated as gel, lotion, cream, oil-in-water emulsion, water-in-oil emulsion and the like.

Preferably the composition is in the form of a water-in-oil emulsion comprising a continuous oil phase and a discontinuous aqueous phase.

The oil phase can comprise silicone oils, non-silicone organic oils, or mixtures thereof, in addition to the liquid, polyol carboxylic acid ester.

In preferred embodiments the oil phase comprises a mixture of volatile silicones and non-volatile silicones. The silicones are useful herein for providing skin conditioning properties. The silicone fluid is present in an amount of from about 1% to about 50% by weight. Suitable volatile silicones include cyclic and linear volatile polyorganosiloxanes. The term "nonvolatile" as used herein shall mean the material has a vapour pressure of no more than 0.1 mm Hg at one atmosphere and 25°C. The term "volatile" as used herein shall mean materials which are not nonvolatile or which have a vapour pressure at the same conditions of more than 0.1 mm Hg.

A description of various volatile silicones is found in Todd, et al.. "Volatile Silicone Fluids for Cosmetics", 91 Cosmetics and Toiletries 27-32 (1976).

Preferred cyclic silicones include cyclic dimethyl siloxane chains containing an average of from about 3 to about 9 silicon atoms, preferably from about 4 to about 5 silicon atoms. Preferred linear silicones include the polydimethylsiloxanes containing an average of from about 3 to about 9 silicon atoms. The linear volatile silicones generally have viscosities of less than about 5 centistokes at 25°C, while the cyclic materials have viscosities of less than about 10 centistokes. Examples of silicone oils useful in the present invention include: Dow Corning 344, Dow Corning 21330, Dow Corning 345, and Dow Corning 200 (manufactured by the Dow Corning Corporation): Silicone 7207 and Silicone 7158 (manufactured by the Union Carbide Corporation). SF:202 (manufactured by General Electric) and SWS-03314 (manufactured by Stauffer Chemical).

The nonvolatile silicones will have vapour pressures as previously defined, and preferably will have an average viscosity of from about 10 to about 100,000 cps at 25°C, more preferably from about 100 to about 10,000 cps, even more preferably from about 500 to about 6000 cps.

9

Lower viscosity non-volatile silicone conditioning agents, however, can also be used. Viscosity can be measured by means of a glass capillary viscometer as set forth in Dow Corning Corporate Test Method CTM0004, July 20, 1970.

Suitable non-volatile silicone fluids for use herein include polyalkyl siloxanes, polyaryl siloxanes, polyalkylaryl siloxanes, polysiloxanes with amino functional substitutions, polyether siloxane copolymers, and mixtures thereof. The siloxanes useful in the present invention may be substituted and/or endcapped with any number of moieties, so long as the material remains suitable for use in a topical cosmetic product, including, for example, methyl, hydroxyl, ethylene oxide, propylene oxide, amino and carboxyl. However, other silicone fluids having skin conditioning properties may be used. The non-volatile polyalkyl siloxane fluids that may be used include, for example, polydimethylsiloxanes. These siloxanes are available, for example, from the General Electric Company as a Viscasil (RTM) series and from Dow Corning as the Dow Corning 200 series. Preferably, the viscosity ranges from about 10 mm².s⁻¹ to about 100,000 mm².s⁻¹ at 25°C. The polyalkylaryl siloxane fluids that may be used, also include, for example, polymethylphenylsiloxanes. These siloxanes are available, for example, from the General Electric Company as SF 1075 methyl phenyl fluid or from Dow Corning as 556 Cosmetic Grade Fluid. The polyether siloxane copolymer that may be used includes, for example, a polypropylene oxide modified dimethylpolysiloxane (e.g., Dow Corning DC-1248) although ethylene oxide or mixtures of ethylene oxide and propylene oxide may also be used.

References disclosing suitable silicone fluids include US-A-2,826,551, Green; US-A-3,964,500, Drakoff, issued June 22nd, 1976; US-A-4,364,837, Pader; and GB-A-849,433, Woolston. In addition, Silicone Compounds distributed by Petrarch Systems Inc., 1984 provides an extensive (though not exclusive) listing of suitable silicone fluids.

Preferred non-volatile silicones for use herein include polydiorganosiloxane-polyoxyalkylene copolymers containing at least one polydiorganosiloxane segment and at least one polyoxyalkylene segment. The polydiorganosiloxane segment has the general formula:

$$R_bSiO_{(4-b)/2}$$

siloxane units wherein b has a value of from about 0 to about 3, inclusive, there being an average value of approximately two R radicals per silicon for all siloxane units in the copolymer, and R denotes a radical selected from methyl, ethyl, vinyl, phenyl and a divalent radical bonding said polyoxyalkylene segment to the polydiorganosiloxane segment. The polyoxyalkylene segment has an average molecular weight of at least about 500, preferably at least about 1000, and comprising from about 0 to about 50 mol percent polyoxypropylene units and from about 50 to about 100 mol percent polyoxyethylene units, at least one terminal portion of said polyoxyalkylene segment being grafted to, or covalently bonded directly or indirectly to a polydiorganosiloxane segment, any terminal portion of said polyoxyalkylene segment not bonded to said polydiorganosiloxane segment being satisfied by a terminating radical; the weight ratio of polydiorganosiloxane segments to polyoxyalkylene segments in said copolymer preferably having a value of from about 2 to about 8. Such polymers are described in US-A-4,268,499.

Preferred for use herein are polydiorganosiloxane-polyoxyalkylene copolymers having the general formula:

$$R^{1} - S_{i} - O = \begin{cases} R^{1} \\ S_{i} - O \\ R^{1} \end{cases} \begin{cases} R^{1} \\ S_{i} - O \\ (CH_{2})_{z} \end{cases} \begin{cases} R^{1} \\ S_{i} - R^{1} \\ R^{1} \end{cases}$$

$$(C_{2}H_{4}O)_{a}(C_{3}H_{6}O)_{b}R$$

wherein R¹ is selected from C1 to C5 alkyl groups, preferably methyl, z is in the range of from 1 to 4, x and y are selected such that the weight ratio of polydiorganosiloxane segments to polyoxalkylene segments is from about 2 to about 8, the mol ratio of a:(a+b) is from about 0.5 to about 1, and R is a chain terminating group, especially selected from hydrogen; hydroxyl; alkyl, such as methyl, ethyl, propyl, butyl, benzyl; aryl, such as phenyl; alkoxy such as methoxy, ethoxy, propoxy, butoxy; benzyloxy; aryloxy, such as phenoxy; alkenyloxy, such as vinyloxy and allyloxy; acyloxy, such as acetoxy, acryloxy and propionoxy and amino, such as dimethylamino.

More preferred for use herein are polydiorganosiloxane-polyoxyalkylene copolymers having the formula:

$$^{\text{CH}_3}$$
 $^{\text{CH}_3}$ $^{\text$

wherein x, y and R are as defined above.

The number of and average molecular weights of the segments in the copolymer are such that the weight ratio of polydiorganosiloxane segments to polyoxyalkylene segments in the copolymer is preferably from about 2.5 to about 4.0.

Suitable copolymers are available commercially under the tradenames Belsil (RTM) from Wacker-Chemie GmbH, Geschaftsbereich S, Postfach D-8000 Munich 22 and Abil (RTM) from Th. Goldschmidt Ltd,. Tego House, Victoria Road, Ruislip, Middlesex, HA4 0YL. Particularly preferred for use herein are Belsil (RTM) 6031, Abil (RTM) B88183,

DC3225C, DC5200, Abil We09 and Abil EM90. A preferred silicone herein is known by its CTFA designation as dimethicone copolyol.

The compositions of the present invention preferably comprise from about 20% to about 95% by weight of composition of oil phase. The oil phase preferably comprises from about 0.01% to about 25%, more preferably from about 0.05% to about 15% by weight of the oil phase of non-volatile silicones. The oil phase preferably comprises from about 35% to about 95%, more preferably from about 45% to about 75% by weight of the oil phase of volatile silicones.

The oil phase in the water-in-oil emulsions herein can also comprise one or more non-silicone organic oil, in addition to the polyol carboxylic acid ester, such as natural or synthetic oil selected from mineral, vegetable, and animal oils, fats and waxes, fatty acid esters, fatty alcohols, fatty acids and mixtures thereof, which ingredients are useful for achieving emollient cosmetic properties. It will be understood that the oil phase may contain, for example, up to about 15%, preferably up to only about 5% of oil phase soluble emulsifier ingredients. Such ingredients are not to be considered as oil phase components from the viewpoint of determining the oil phase level.

Suitable organic oils for use herein include, for example, optionally hydroxy-substituted C8-C50 unsaturated fatty acids and esters thereof, C1-C24 esters of C8-C30 saturated fatty acids such as isopropyl myristate, isopropyl palmitate, cetyl palmitate and octyldodecylmyristate (Wickenol 142), beeswax, saturated and unsaturated fatty alcohols such as behenyl alcohol and cetyl alcohol, hydrocarbons such as mineral oils, petrolatum and squalane, fatty sorbitan esters (see US-A-3988255, Seiden, issued October 26 1976), lanolin and lanolin derivatives, animal and vegetable triglycerides such as almond oil, peanut oil, wheat germ oil, linseed oil, jojoba oil, oil of apricot pits, walnuts, palm nuts, pistachio nuts, sesame seeds, rapeseed, cade oil, corn oil, peach pit oil, poppyseed oil, pine oil, castor oil, soybean oil, avocado oil, safflower oil, coconut oil, hazelnut oil, olive oil, grapeseed oil, shea butter, shorea butter, and sunflower seed oil and C1-C24 esters of dimer and trimer acids such as diisopropyl dimerate,

diisostearylmalate, diisostearyldimerate and triisostearyltrimerate. Of the above, highly preferred are the mineral oils, petrolatums, unsaturated fatty acids and esters thereof and mixtures thereof.

The compositions herein preferably further comprise water at a level of from about 10% to about 60%, preferably from about 30% to about 50% by weight of composition.

Acidic Skin Care Active

The compositions of the present invention can comprise an acidic skin care active.

Suitable acidic skin care actives can be selected from hydroxycarboxylic acids. As used herein the term acidic skin care active means any skin care active containing an acidic functional group (e.g. carboxy, sulfonic).

Suitable hydroxycarboxylic acids can be selected from hydroxymonocarboxylic acids having the following chemical structure:

R₁(CR₂OH)_m(CH₂)_nCOOH

wherein R_1 , $R_2 = H$, alkyl, aralkyl or aryl group of saturated or unsaturated, straight or branched chain or cyclic form, having from 1 to 25 carbon atoms; m=1,2,3,4,5,6,7,8 or 9; n=0 or a numerical number up to 23.

The hydroxymonocarboxylic acid may be present as a free acid, lactone, or salt form, The lactone form could be either inter or intramolecular lactone, however, most common ones are intramolecular lactones with a ring structure formed by elimination of one or more water molecules between a hydroxy group and the carboxylic group. Since the hydroxymonocarboxylic acids are organic in nature, they may form a salt or a complex with an inorganic or organic base such as ammonium hydroxide, sodium or potassium hydroxide, or triethanolamine.

The hydroxymonocarboxylic acid and its related compounds may exist as stereoisomers such as D, L, and DL forms.

Typical alkyl, aralkyl and aryl groups for R1 and R2 include methyl, ethyl, propyl, isopropyl, benzyl and phenyl. The hydrogen atoms of the R₁ and R₂ and (CH₂)_n may be substituted by a nonfunctional element such as F, Cl, Br, I, S or a radical such as a lower alkyl or alkoxy, saturated or unsaturated, having 1 to 9 carbon atoms. Representative hydroxymonocarboxylic acids are 2-hydroxyacetic acid (glycolic acid), 2hydroxypropanoic acid (lactic acid), 2-methyl 2-hydroxypropanoic acid (methyllactic acid), 2-hydroxybutanoic acid, phenyl 2-hydroxyacetic acid (mandelic acid), phenyl 2-methyl 2-hydroxyacetic acid, 3-phenyl 2hydroxypropanoic acid (phenyllactic acid), 2,3-dihydroxypropanoic acid (glyceric acid), 2.3.4-trihydroxybutanoic acid, 2,3,4,5tetrahydroxypentanoic acid, 2,3,4,5,6-pentahydroxyhexanoic acid, 2hydroxydodecanoic acid (alpha hydroxylauric acid), 2,3,4,5,6,7hexahydroxyheptanoic acid, diphenyl 2-hydroxyacetic acid (benzilic acid), 4-hydroxymandelic acid, 4-chloromandelic acid, 3-hydroxybutanoic acid, 4-hydroxybutanoic acid, 2-hydroxyhexanoic acid, 5-hydroxydodecanoic acid, 12-hydroxydodecanoic acid, 10-hydroxydecanoic acid, 16hydroxyhexadecanoic acid, 2-hydroxy-3-methylbutanoic acid, 2-hydroxy-4-methylpentanoic acid, 3-hydroxy-4-methoxymandelic acid, 4-hydroxy-3-methoxymendelic acid, 2-hydroxy-2-methylbutanoic acid, 3-(2hydroxyphenyl) lactic acid, 3-(4-hydroxyphenyl) lactic acid, hexahydromandelic acid, 3-hydroxy-3-methylpentanoic acid, 4hydroxydecanoic acid, 5-hydroxydecanoic acid and aleuritic acid.

Another type of hydroxyacid suitable for use herein is a hydroxydicarboxylic acid having the following formula:

HOOC(CHOH)_m(CH₂)_nCOOH

wherein m=1,2,3,4,5,6,7,8 or 9; n=0 or an integer up to 23.

The hydroxydicarboxylic acid may also be present as a free acid, lactone or salt form. The hydroxydicarboxylic acid and its related compounds may also exist as stereoisomers such as D, L, DL and meso forms.

The hydrogen attached to the carbon atom may be substituted by a nonfunctional element such as F, Cl, Br, I, S, or a radical such as a lower saturated or unsaturated alkyl or alkoxy having from 1 to 9 carbon atoms.

Representative hydroxydicarboxylic acids are 2-hydroxypropanedioic acid (tartronic acid), 2-hydroxybutanedioic acid (malic acid), erythraric acid and threaric acid (tartaric acid), arabiraric acid, ribaric acid, xylaric acid and lyxaric acid, glucaric acid (saccharic acid), galactaric acid (mucic acid), mannaric acid, gularic acid, allaric acid, altraric acid, idaric acid and talaric acid.

A third type of hydroxyacid suitable for use herein is a miscellaneous group of compounds which is not readily represented by the above generic structure of either the first type or the second type described above. Included in the third type of hydroxyacids are the following:

Hydroxycarboxylic acids of formula:

$R(OH)_m(COOH)_n$

wherein m, n=1,2,3,4,5,6,7,8 or 9, R=H, alkyl, aralkyl or aryl group of saturated or unsaturated, straight or branched chain or cyclic form, having from 1 to 25 carbon atoms; citric acid, isocitric acid, citramalic acid, agaricic acid (n-hexadecylcitric acid), quinic acid, uronic acids including glucuronic acid, glucuronolactone, galacturonic acid, galacturonolactone, hydroxypyruvic acid, hydroxypyruvic acid phosphate, ascorbic acid, dihydroxypyruvic acid, 2-hydroxy-2-methylbutanoic acid, 1-hydroxy-1-cyclopropane carboxylic acid, 2-hydroxyhexanedial, 5-hydroxylysine, 3-hydroxy-2-aminopentanoic acid, tropic acid, 4-hydroxy-2,2-diphenylbutanoic acid, 3-hydroxy-3-methylglutaric acid, and 4-hydroxy-3-pentenoic acid.

The third type of hydroxyacid may also be present as a free acid, lactone or salt form and may also exist as stereoisomers such as D, L, DL and meso forms.

The hydrogen atom attached to the carbon atom may be substituted by a nonfunctional element such as F, Cl, Br, I, S or a radical such as a lower saturated or unsaturated alkyl or alkoxy having from 1 to 9 carbon atoms.

Mixtures of hydroxy acids can also be used in the compositions herein. Hydroxy acids are useful herein from the viewpoint of reducing wrinkles and improving skin feel and appearance.

Other suitable hydroxy acids for use herein include salicylic acid, retinoic acid, and azelaic acid.

Preferred acidic skin care actives for use herein include salicylic acid, retinoic acid, azelaic acid, lactic acid, glycolic acid, pyruvic acid, and mixtures thereof, more preferably salicylic acid and azelaic acid, and especially salicylic acid. The salicylic acid is useful herein as an anti-acne active.

Preferably, the acidic skin care active is present at a level of from about 0.1% to about 10%, preferably from about 0.1% to about 5%, more preferably from about 0.5% to about 3%, by weight of composition.

The acidic skin care active is solubilized in water or a hydroalcoholic solution, for example, solutions based upon C_2 - C_6 alcohols, diols and polyols, preferred alcohols being selected from ethanol, dipropylene glycol, butylene glycol, hexylene glycol, and mixtures thereof. Alcohol is preferably present in the compositions herein at a level of from about 1% to about 20%. The final aqueous/hydroalcoholic skin care active solution preferably has a pH at ambient temperature (25°C) of less than about p K_a +1, where p K_a is the logarithmic acidity constant for the fully protonated skin care active. In preferred embodiments, the pH of the final solution is less than about p K_a .

The logarithmic acidity constant is thus defined by reference to the equilibrium

$$H_{n}A = H^{+} + H_{n-1}A^{-}$$

where H_nA is the fully protonated acid, n is the number of protons in the fully protonated acid and H_{n-1} A is the conjugate base of the acid corresponding to loss of one proton.

The acidity constant for this equilibrium is therefore

$$K_n = \underline{[H^+][H_{n-1}A]}$$
$$[H_nA]$$

and
$$pK_a = -\log_{10}K_n$$

For the purposes of this specification, acidity constants are defined at 25°C and at zero ionic strength. Literature values are taken where possible (see Stability Constants of Metal-Ion Complexes, Special Publication No. 25, The Chemical Society, London); where doubt arises they are determined by potentiometric titration using a glass electrode.

The pK_a of the acidic skin care active used herein is preferably in the range of from about 1 to about 5.5, more preferably from about 2 to about 4.5, especially from about 2 to about 4.

When the composition contains an acidic skin care active, the pH of the aqueous phase is preferably less than about pH 6, more preferably from about pH 2 to about pH 5, especially from about pH 2.5 to about pH 4. At pH values of less than about 5 the aqueous phase is preferably free of acid labile species such as polyacrylic or polymethacrylic acids or esters.

The compositions of the present invention can also comprise a solubilizing agent for solubilizing the acidic skin care active. Any solublizing agent suitable for use in a cosmetic composition can be used. Preferably the solubilizing agent herein is selected from polyoxyethylene-

polyoxypropylene ethers of C4 to C22 alcohols, pyrrolidone-based solubilising agents, polyethylene glycol based nonionic surfactants having an HLB of greater than about 15, preferably greater than about 18, and mixtures thereof.

Pyrrolidone-based solubilising agents suitable for use herein include polyvinylpyrrolidone or C1-C4 alkyl polyvinylpyrrolidone having a molecular weight (viscosity average) in the range from about 1500 to about 1,500,000, preferably from about 3000 to about 700,000, more preferably from about 5000 to about 100,000. Suitable examples of pyrrolidone-based solubilising agents are polyvinylpyrrolidone (PVP) (or povidone) and butylated polyvinylpyrrolidone. The most preferred pyrrolidone-based solubilising agent herein is polyvinylpyrrolidone. PVP is commercially available under the trade name Luviskol (RTM) from BASF. A preferred PVP solubilising agent herein is Luviskol K17 which has a viscosity-average molecular weight of about 9,000. Other pyrrolidone-based solubilising agents for use herein include C1-C18 alkyl or hydroxyalkyl pyrrolidones such as lauryl pyrrolidone.

The pyrrolidone-based solubilising agent is preferably present in the composition herein in a level of from about 0.1% to about 10%, more preferably from about 0.1% to about 5%, especially from about 0.5% to about 2% by weight of composition. The weight ratio of acidic skin care active: pyrrolidone-based solubilising agent is preferably in the range from about 10:1 to about 1:10, more preferably from about 5:1 to about 1:5.

Preferred embodiments of the invention additionally comprise from about 0.01% to about 5% by weight of an additional acid or salt thereof which is soluble in water at pH values of less than or equal to the pKa of the corresponding acid, for example, an acid selected from citric acid, boric acid, and salts, and mixtures thereof. These materials are valuable herein in combination with the pyrrolidone-based complexing agent from the viewpoint of aiding solubilization of the acidic skin care active. Particularly preferred herein from this viewpoint is a sodium salt of citric

acid. In preferred embodiments, the acid or salt thereof is soluble to a level of at least 5% w/w at 25°C.

A preferred solubilizing agent in the compositions of the present invention is a nonionic surfactant selected from polyoxyethylene-polyoxypropylene ethers of C4-C22 alcohols, and mixtures thereof. The nonionic surfactant is valuable herein as a solubilising agent for the acidic skin care active in the discontinuous aqueous phase. Suitable polyoxyethylene-polyoxypropylene ethers of C4-C22 alcohols for use herein include those having the general formula:

R(OCHCH₂)_x(OCH₂CH₂)_yOH CH₃

wherein x is in the range of from about 1 to about 35, preferably from about 1 to about 10, y is in the range of from about 1 to about 45, preferably from about 1 to about 30 and R is a straight chain or branched chain C4 to C22 alkyl group, or a mixture thereof. In preferred embodiments (x+y) is greater than or equal to 5, preferably greater than or equal 10, more preferably greater than or equal to 15. The ratio of x:y is in the range from 1:1 to 1:10.

Examples of suitable R groups in the above formula include cetyl, butyl, stearyl, cetearyl, decyl, lauryl and myristyl.

Examples of suitable polyoxyethylene-polyoxypropylene alcohol ethers include (using CTFA designations) PPG-4-Ceteth-1, PPG-4-Ceteth-5, PPG-4-Ceteth-10, PPG-4-Ceteth-20, PPG-5-Ceteth-20, PPG-8-Ceteth-1, PPG-8-Ceteth-2, PPG-8-Ceteth-5, PPG-8-Ceteth-10, PPG-8-Ceteth-20, PPG-2-Buteth-3, PPG-2-Buteth-5, PPG-5-Buteth-7, PPG-9-Buteth-12, PPG-28-Buteth-35, PPG-12-Buteth-16, PPG-15-Buteth-20, PPG-20-Buteth-30, PPG-24-Buteth-27, PPG-26-Buteth-26, PPG-33-Buteth-45, PPG-2-Ceteareth-9, PPG-4-Ceteareth-12, PPG-10-Ceteareth-20, PPG-2-Deceth-10, PPG-4-Deceth-4, PPG-6-Deceth-4, PPG-6-Deceth-9, PPG-8-Deceth-6, PPG-2-Isodeceth-4, PPG-2-Isodeceth-9, PPG-2-Isodeceth-9, PPG-2-Isodeceth-12, PPG-3-Isodeceth-1, PPG-4-Laureth-5, PPG-4-

Laureth-2, PPG-4-Laureth-7, PPG-5-Laureth-5, PPG-25-Laureth-25, PPG-3-Myreth-11, PPG-3-Myreth-3 and PPG-9-Steareth-3.

Preferred polyoxyethylene-polyoxypropylene ethers for use herein are ethers of C8 to C16 alcohols having the formula (I) wherein x is from 2 to 12 and y is from 10 to 30 and where the ratio of x:y is in the range of from about 1:2 to about 1:8.

Particularly preferred polyoxyethylene-polyoxypropylene ethers of C4 to C22 alcohols for use herein are those having the formula (I) above wherein R is cetyl and wherein x is in the range of from about 4 to about 8, and wherein y is in the range of from about 15 to about 25, and the ratio of x:y is in the range of from about 1:3 to about 1:5. A particularly preferred ether from the viewpoint of improving solubilisation of the acidic skin care active is PPG-5-Ceteth-20, which is available under the tradename Procetyl AWS.

The solubilizing agent herein is preferably present at a level of from about 0.1% to about 15%, more preferably from about 1% to about 10%, especially from about 2% to about 8% by weight of composition.

Preferred embodiments herein comprise a pigment or mixture of pigments. The pigment used herein must be compatible with any acidic skin care active which is present in the composition and have excellent overall colour stability. Suitable pigments for use herein can be inorganic and/or organic. Also included within the term pigment are materials having a low colour or lustre such as matte finishing agents, and also light scattering agents. Examples of suitable pigments are iron oxides, rutile titanium dioxide, anatase titanium dioxide, ferric oxide, ferrous oxide, chromium oxide, chromium hydroxide, manganese violet, acylglutamate iron oxides, ultramarine blue, D&C dyes, carmine, and mixtures thereof. Depending upon the type of make-up composition, eg. foundation or blusher, a mixture of pigments will normally be used.

The foundation composition can also include at least one matte finishing agent. The function of the matte finishing agent is to hide skin defects and

reduce shine. Such cosmetically acceptable inorganic agents, i.e., those included in the CTFA Cosmetic Ingredient Dictionary, Third Ed., as silica, hydrated silica, silicone-treated silica beads, mica, talc, polyethylene, titanium dioxide, bentonite, hectorite, kaolin, chalk, diatomaceous earth, attapulgite zinc oxide and the like may be utilized. Of particular usefulness as a matte finishing agent is low lustre pigment such as titanated mica (mica coated with titanium dioxide) coated with barium sulfate. Of the inorganic components useful as a matte finishing agent low lustre pigment, talc, polyethylene, hydrated silica, kaolin, titanium dioxide and mixtures thereof are particularly preferred. Materials suitable for use herein as light-scattering agents can be generally described as spherical shaped inorganic materials having a particle size of up to about 100 microns, preferably from about 5 to about 50 microns, for example spherical silica particles.

Other examples of pigments include lakes of organic colorants such as FD&C Red No. 7 calcium lake, FD&C Yellow No. 5 aluminium lake, D&C Red No. 9 barium lake, and D&C Red No. 30.

The preferred pigments for use herein from the viewpoint of moisturisation, skin feel, skin appearance and emulsion compatibility are treated pigments. The pigments can be treated with compounds such as amino acids such as lysine, silicones, lauroyl, collagen, polyethylene, lecithin and ester oils. The more preferred pigments are the silicone (polysiloxane) treated pigments.

A highly preferred pigment for use herein is a pigment which has been coated with organosilicon component selected from a polyorganosiloxane or a silane wherein the coated pigment has a hydrogen potential of less than about 2.0, preferably less than about 1.0, more preferably less than about 0.5 ml, and especially less than about 0.1 ml H2/g of coated pigment. The pigment preferred for use herein is in particulate form. The pigment is incorporated into the continuous oil phase in the compositions herein. The coatings used can be bonded to the pigment surface by covalent bonding, physical adsorption or adhesion, preferably by covalent bonding to the surface of the pigment. The function of the coatings herein is to

hydrophobically-modify the pigments so that thay are "wettable" in the continuous silicone phase of the water-in-silicone emulsions. The coated pigment is also useful herein from the viewpoint of reducing hydrogen gas evolution and improving product stability.

Without wishing to be limited by theory it is believed that although the pigments are present in the oil phase of the water-in-oil emulsion, hydrogen ions from the aqueous phase can pass through the interface of the emulsion into the oil phase, where they are available to react with the pigment coatings, e.g. to give off hydrogen gas. However, by using organosilicon-coated pigments having a hydrogen potential of less than about 2 ml H₂/g of coated pigment, hydrogen gas generation is reduced.

The hydrogen potential of the coated pigment is measured herein using the following test method:

A dispersion of the coated pigment containing 20g of coated pigment is placed in a flask on a magnetic stirrer and 100ml of a 2% ethanolic solution of potassium hydroxide is added with stirring at ambient temperature. The hydrogen gas which is evolved is collected in a second flask at ambient temperature and pressure (25°C, 1At). The hydrogen gas released can therefore be volumetrically measured.

A wide variety of organosilicon components can be used for treating the pigments herein. A suitable polyorganosiloxane herein is selected from:

(A) material of the formula:

$$(R^1)_3$$
SiO- $(Si(R^2R^3)O)_p$ -Si $(R^2R^3)OA_2$

wherein p is 1 to 1000, preferably from 1 to 100, A_2 is hydrogen or an alkyl group having from 1 to 30 carbon atoms, R^1 is a C_1 - C_{30} alkyl, preferably methyl, R^2 and R^3 are independently selected from a C_1 - C_{30} alkyl and a phenyl, preferably wherein R^2 and R^3 are both methyl or wherein R^2 is methyl and R^3 is phenyl; or

(B) material of the formula:

$$(R^1)_3 SiO(Si(R^2)(H)-O)_i-Si(R^1)_3$$

wherein i is 1 to 1000, preferably 1 to 100, and wherein R1 and R2 are as defined above for formula (A).

In preferred embodiments the organosilicon component is selected from a silane. The silane can be selected from material of the formula:

(C) $A_1SiX_1X_2X_3$

wherein A is an alkyl or alkenyl group having from 1 to 30 carbon atoms, and X_1 , X_2 and X_3 are independently C_1 - C_4 alkoxy preferably methoxy or ethoxy, or halo, preferably chloro.

When the pigment herein is treated with silane material having the formula (C) described herein above a pigment having the following formula (1) is produced:

$$P-O-Si(OH)(A)-[-O-Si(OH)(A)-]_{0-100}-OH$$

wherein P is an atom in the pigment surface and each A is an alkyl or alkenyl group having up to 30 carbon atoms. A number of adjacent polysiloxane chains as shown in formula (1) can be cross-linked through oxygen atoms to form a polysiloxane chain with up to 100 repeating -Si(-OP)-O-units that extend along the pigment surface, in addition to the polysiloxane chain which extends away from the pigment surface. Examples of linear or branched alkyl groups are methyl, ethyl, propyl, butyl, pentyl, hexyl, heptyl, octyl, and so forth up to octadecyl. "Alkenyl" includes carbon chains with one or more double bonds; examples of such groups include ethylene, propylene, acrylyl, methacrylyl, and residues of unsaturated fatty acids such as oleic (C₁₇C₃₃-), linoleic (C₁₇H₃₁-), and linolenic (C₁₇H₂₉-).

When the pigment herein is treated with polyorganosiloxane material having the formula (A) described hereinabove a pigment having the following formula (2) is produced:

P-O-
$$(Si(R^2R^3)O)_p$$
- $Si(R^1)_3$ (2)

wherein p is 1-1000, preferably 1 to 100, R¹, R² and R³ are as defined above for formula (A) and P is an atom in the pigment surface.

When the pigment herein is treated with polyorganosiloxane material having the formula (B) described hereinabove a pigment having the following formula (3) is produced:

$$(R^1)_3SiO-[Si(R^2)(-OP)-O-]_p-Si(R^1)_3$$
 (3)

wherein each P is an atom in the pigment surface, p is from 1 to 1000, preferably from 1 to 100, R¹ and R² are as defined above in formula (B) and in which each of the up to 100 repeating (Si-O) units is bonded through an oxygen atom to the pigment surface.

The pigment (or a mixture of two or more pigments) can be coated by placing it in dry, finely divided form in a mixer, adding the organosilicon component, and mixing. The organosilicon coating is preferably present at a level of from about 0.01% to about 5%, more preferably from about 0.1% to about 4%, and especially from about 0.5% to about 2%, by weight of the organosilicon coated pigment.

The most preferred coated pigment from the viewpoint of reducing hydrogen gas evolution and improving product stability is Cardre 70429.

The total concentration of the coated pigment may be from about 0.1 to about 25% by weight and is preferably from about 1 to about 15%, more preferably from about 8% to about 12% by weight of the total composition, the exact concentration being dependent to some extent upon the specific mixture of pigments selected for use in a foundation make-up or blusher to achieve the desired shades. The preferred compositions

contain from about 2% to about 20% by weight of titanium dioxide and most preferably from about 5% to about 10% by weight of titanium dioxide.

A highly preferred component of the compositions herein is a humectant or mixture of humectants. The humectant or mixture of humectants herein is present in an amount of from about 0.1% to about 30% preferably from about 1% to about 25%, and more preferably from about 1% to about 10% by weight of composition. Suitable humectants are selected from glycerine and polyglycerylmethacrylate lubricant having a viscosity at 25°C of 300,000 to 1,100,000 cps; a specific gravity at 25°C of 1 to 1.2g/ml, a pH of 5.0 to 5.5; a bound water content of 33 to 58%; and, a free water content from 5 to 20%.

The humectant can be incorporated at least partly into the oil phase of the water-in-oil emulsion. The oil phase preferably comprises from about 0.1% to about 10%, more preferably from about 0.1% to about 3% by weight of humectant on a composition basis. The humectant can be introduced into the oil phase in the form of a mixture with or incorporated within a particulate lipophilic or hydrophobic carrier material.

Polyglycerylmethacrylate lubricants having the desired properties are marketed by Guardian Chemical Corporation under the trademark "Lubrajel". The "Lubrajels" identified as "Lubrajel DV", "Lubrajel MS", and "Lubrajel CG" are preferred in the present invention. The gelling agents sold under these trademarks contain about 1% propylene glycol.

Other suitable humectants include sorbitol, panthenols, propylene glycol, dipropylene glycol, butylene glycol, hexylene glycol, alkoxylated glucose derivatives, such as Glucam (RTM) E-20, hexanetriol, and glucose ethers, and mixtures thereof.

The panthenol moisturiser can be selected from D-panthenol ([R]-2,4-dihydroxy-N-[3-hydroxypropyl)]-3,3-dimethylbutamide), DL-panthenol, calcium pantothenate, royal jelly, panthetine, pantotheine, panthenyl ethyl ether, pangamic acid, pyridoxin, pantoyl lactose and Vitamin B complex.

The preferred humectant herein is glycerine. Chemically, glycerine is 1,2,3-propanetriol and is a product of commerce.

The make-up compositions of the present invention can also comprise a particulate cross-linked hydrophobic acrylate or methacrylate copolymer. This copolymer is particularly valuable for reducing shine and controlling oil while helping to provide effective moisturization benefits. The cross-linked hydrophobic polymer is preferably in the form of a copolymer lattice with at least one active ingredient dispersed uniformly throughout and entrapped within the copolymer lattice. Alternatively, the hydrophobic polymer can take the form of a porous particle having a surface area (N2-BET) in the range from about 50 to 500, preferably 100 to 300m²/g and having the active ingredient absorbed therein.

The cross-linked hydrophobic polymer when used herein is in an amount of from about 0.1% to about 10% by weight and is preferably incorporated in the external silicone-containing oil phase. The active ingredient can be one or more or a mixture of skin compatible oils, skin compatible humectants, emollients, moisturizing agents and sunscreens. The polymer material is in the form of a powder, the powder being a combined system of particles. The system of powder particles forms a lattice which includes unit particles of less than about one micron in average diameter, agglomerates of fused unit particles of sizes in the range of about 20 to 100 microns in average diameter and aggregates of clusters of fused agglomerates of sizes in the range of about 200 to 1,200 microns in average diameter.

The powder material of the present invention which can be employed as the carrier for the active ingredient can be broadly described as a cross-linked "post absorbed" hydrophobic polymer lattice. The powder preferably has entrapped and dispersed therein, an active which may be in the form of a solid, liquid or gas. The lattice is in particulate form and constitutes free flowing discrete solid particles when loaded with the active material. The lattice may contain a predetermined quantity of the active material. The polymer has the structural formula:

where the ratio of x to y is 80:20, R' is -CH₂CH₂- and R" is - $(CH_2)_{11}CH_3$.

A suitable hydrophobic polymer for use herein is a highly crosslinked polymer, more particularly a highly cross-linked polymethacrylate copolymer such as that manufactured by the Dow Corning Corporation, Midland. Michigan, USA, and sold under the trademark POLYTRAP (RTM). It is an ultralight free-flowing white powder and the particles are capable of absorbing high levels of lipophilic liquids and some hydrophilic liquids while at the same time maintaining a free-flowing powder character. The powder structure consists of a lattice of unit particles less than one micron that are fused into agglomerates of 20 to 100 microns and the agglomerates are loosely clustered into macroparticles or aggregates of about 200 to about 1200 micron size. The polymer powder is capable of containing as much as four times its weight of fluids, emulsions, dispersions or melted solids.

Adsorption of actives onto the polymer powder can be accomplished using a stainless steel mixing bowl and a spoon, wherein the active is added to the powder and the spoon is used to gently fold the active into the polymer powder. Low viscosity fluids may be adsorbed by addition of the fluids to a sealable vessel containing the polymer and then tumbling the materials until a consistency is achieved. More elaborate blending equipment such

as ribbon or twin cone blenders can also be employed. The preferred active ingredient for use herein is glycerine. Preferably, the weight ratio of humectant: carrier is from about 1:4 to about 3:1.

Also suitable as a highly cross-linked polymethacrylate copolymer is Microsponge 5640. This takes the form of generally spherical particles of cross-linked hydrophobic polymer having a pore size of from about 0.01 to about 0.05 µm and a surface area of 200-300m²/g. Again, it is preferably loaded with humectant in the levels described above.

The compositions of the invention can also contain a hydrophilic gelling agent at a level preferably from about 0.01% to about 10%, more preferably from about 0.02% to about 2%, and especially from about 0.02% to about 0.5%. The gelling agent preferably has a viscosity (1% aqueous solution, 20°C, Brookfield RVT) of at least about 4000 mPa.s, more preferably at least about 10,000 mPa.s and especially at least 50,000 mPa.s.

Suitable hydrophilic gelling agents can generally be described as water-soluble or colloidally water-soluble polymers, and include cellulose ethers (e.g. hydroxyethyl cellulose, methyl cellulose, hydroxypropylmethyl cellulose), polyvinylalcohol, polyquaternium-10, guar gum, hydroxypropyl guar gum and xanthan gum.

Among suitable hydrophilic gelling agents are acrylic acid/alkyl acrylate copolymers and the carboxyvinyl polymers sold by the B.F. Goodrich Company under the trade mark of Carbopol resins. These resins consist essentially of a colloidally water-soluble polyalkenyl polyether crosslinked polymer of acrylic acid crosslinked with from 0.75% to 2.00% of a crosslinking agent such as for example polyallyl sucrose or polyallyl pentaerythritol. Examples include Carbopol 934, Carbopol 940, Carbopol 950, Carbopol 980, Carbopol 951 and Carbopol 981. Carbopol 934 is a water-soluble polymer of acrylic acid crosslinked with about 1% of a polyallyl ether of sucrose having an average of about 5.8 allyl groups for each sucrose molecule. Also suitable for use herein are hydrophobically-modified cross-linked polymers of acrylic acid having amphipathic

properties available under the Trade Name Carbopol 1382, Carbopol 1342 and Pemulen TR-1 (CTFA Designation: Acrylates/10-30 Alkyl Acrylate Crosspolymer). A combination of the polyalkenyl polyether cross-linked acrylic acid polymer and the hydrophobically modified cross-linked acrylic acid polymer is also suitable for use herein. Other suitable gelling agents suitable for use herein are oleogels such as trihydroxystearin and aluminium magnesium hydroxy stearate. The gelling agents herein are particularly valuable for providing excellent stability characteristics over both normal and elevated temperatures.

Preferably the acidic group containing hydrophilic gelling agents are neutralized. Neutralizing agents suitable for use in neutralizing acidic group containing hydrophilic gelling agents herein include sodium hydroxide, potassium hydroxide, ammonium hydroxide, monoethanolamine, diethanolamine and triethanolamine.

The make-up compositions herein can additionally comprise an emollient. Emollients suitable for the compositions of the present invention include natural and synthetic oils selected from mineral, vegetable, and animal oils, fats and waxes, fatty acid esters, fatty alcohols, alkylene glycol and polyalkylene glycol ethers and esters, fatty acids and mixtures thereof.

Suitable emollients for use herein include, for example, optionally hydroxy-substituted C8-C50 unsaturated fatty acids and esters thereof, C1-C24 esters of C8-C30 saturated fatty acids such as isopropyl myristate, cetyl palmitate and octyldodecylmyristate (Wickenol 142), beeswax, saturated and unsaturated fatty alcohols such as behenyl alcohol and cetyl alcohol, hydrocarbons such as mineral oils, petrolatum and squalane, fatty sorbitan esters (see US-A-3988255, Seiden, issued October 26 1976), lanolin and lanolin derivatives, such as lanolin alcohol ethoxylated, hydroxylated and acetylated lanolins, cholesterol and derivatives thereof, animal and vegetable triglycerides such as almond oil, peanut oil, wheat germ oil, linseed oil, jojoba oil, oil of apricot pits, walnuts, palm nuts, pistachio nuts, sesame seeds, rapeseed, cade oil, corn oil, peach pit oil, poppyseed oil, pine oil, castor oil, soybean oil, avocado oil, safflower oil, coconut oil, hazelnut oil, olive oil, grapeseed oil, and sunflower seed oil

and C₁₋C₂₄ esters of dimer and trimer acids such as diisopropyl dimerate, diisostearylmalate, diisostearyldimerate and triisostearyltrimerate.

Preferred emollients are selected from hydrocarbons such as isohexadecane, mineral oils, petrolatum and squalane, lanolin alcohol, and stearyl alcohol. These emollients may be used independently or in mixtures and may be present in the composition of the present invention in an amount from about 1% to about 30% by weight, and preferably are present in an amount from about 5% to about 15% by weight of the total composition.

The composition may also contain additional materials such as, for example, fragrances, sun-screens, preservatives, electrolytes such as sodium chloride, proteins, antioxidants, chelating agents and water-in-oil emulsifiers as appropriate.

Another optional component of the make-up composition is one or more ultraviolet absorbing agents. Ultraviolet absorbing agents, often described as sunscreening agents, can be present in a concentration in the range of between about 1% and about 12% by weight, based on the total weight of composition. Preferably, the UV absorbing agents constitute between about 2% and 8% by weight. More preferably, the UV absorbing agents can be present in the composition in a concentration range of between about 3% and about 6% by weight. Of the ultraviolet absorbing agents suitable for use herein, benzophenone-3, octyl methocycinnamate, octyl dimethyl PABA (Padimate O), Parsol 1789, Parsol MCX, and mixtures thereof are particularly preferred.

Another optional but preferred component herein is one or more additional chelating agents, preferably in the range of from about 0.02% to about 0.5% by weight, based on the total weight of the composition. Preferably, the chelating agent is present in a concentration in the range of between about 0.03% and about 0.2% by weight, based on the total weight of the composition. Among the chelating agents that may be included in the composition is tetrasodium EDTA.

Another optional but preferred component of the foundation composition is one or more preservatives. The preservative concentration in the foundation composition, based on the total weight of that composition, is in the range of between about 0.05% and about 1% by weight, preferably between about 0.1% and about 0.8% by weight. Suitable preservatives for use herein include sodium benzoate, propyl paraben, phenoxyethanol and mixtures thereof.

The composition preferably comprises from about 20% to about 95%, more preferably from about 30% to about 70% by weight of the oil phase, and from about 5% to about 80%, more preferably from about 30% to about 70% by weight of the aqueous phase. The aqueous phase preferably comprises from about 40% to about 90%, more preferably from about 60% to about 80% by weight of aqueous phase of water.

The cosmetic compositions of the present invention can be in the form of foundations, blushers, concealers, compact powders, and the like, preferably as foundations and concealers.

Method of Use

The compositions herein are used for reducing the appearance of fine lines and wrinkles. The compositions herein can be used by topically applying to the areas of skin to be treated an effective amount of the compositions. As used herein the term "effective amount" means an amount sufficient to reduce the appearance of wrinkles and fine lines.

The composition can be applied for several days, weeks, months or years at appropriate intervals. The compositions are preferably applied from about four times a day to about once every three days, more preferably from about twice a day to once every other day, especially about once a day. Typically, in each application, an effective amount of composition is delivered to the skin by topically applying (in terms of mg/cm² skin) from about 0.01 to about 5 mg composition/cm², more preferably from about 1 to about 2 mg composition/cm² skin.

The compositions are generally applied by lightly massaging the composition into the skin typically in the amounts described above.

The table below shows examples of cosmetic compositions of the present invention.

	1/%	2/%	3/%	4/%	5/%
PHASE A			 	-	<u> </u>
Cyclomethicone Dimethicone copolyol (DC-3225C)1	15	16	15	20	18
Cyclomethicone (DC2-1330)1	6	7.5	7.0	6	8
Cetyl Dimethicone Copolyol	0.2	0.2	0.2	0.25	0.2
Cyclomethicone (DC-200-350cs) ¹	0.5	0	0	1	0.5
Cyclomethicone (DC-200-50cs) ¹	0.5	0	1	2	0
SEFA cottonate ²	4.0	5.0	3.0	4.0	3.0
PHASE B		<u> </u>			
Talc	4	1.2	0	2	3
Mica & Barium Sulphate & TiO2	0.1	0.1	0.1	0	0.5
Silica L-1500 ³	0.5	0.5	1	0.5	1
MT100T (Ti02)4	0.0	0.5	0.5	0	0
Zinc oxide	0.0	0	3	3	0
MT500B (TiO2)4	0	3	0	0	2.0
Dimethicone-coated TiO2 (SAT) ⁵	8.25	6.25	7.0	9	8.25
PHASE C					<u></u>
Black iron oxide pigment grind	0.3	0.45	0.17	0.35	0.32
Red iron oxide pigment grind	1.2	1.33	0.52	0.69	1
Yellow iron oxide pigment grind	2.5	4	1.82	2.38	3.3
PHASE E					
Trihydroxystearin	0.3	0.3	. 0.3	0.3	0.3

Cyclomethicone (DC2-1330) ¹	1.0	1	1	1	1			
PHASE F	<u> </u>							
Laureth-7	0.5	0.5	0.5	0.5	0.5			
Propylparaben	0.25	0.25	0.25	0.25	0.5			
PHASE G								
Nylon-12 (Orgasol 2002 DNAT)6	0.0	5.0	3.0	0.0	3.0			
Nylon-12 (Orgasol 2002 EXDNAT (fine)) ⁶	3.0	0	0	5.0	0.0			
		0.						
PHASE H								
Water	to 100							
Glycerin	10	6	8	10	8			
Polyvinylpyrrolidone (Luviskol K-17) ⁷	0	1	0.5	0	0.5			
Sodium chloride	2.0	2.0	2.0	2.0	2.0			
Sodium dehyroacetate	0.3	0.3	0.3	0.3	0.3			
Phenoxyethanol	0.25	0.25	0.25	0.25	0.25			
Tetra sodium EDTA	0.1	0.1	0.1	0.1	0.1			

- 1. Supplied by Dow Corning Ltd, Avco House, Castle Street, Reading, RG1 7DZ, UK
- 2. Supplied By Procter & Gamble
- 3. Supplied by Presperse Inc., P.O.Box 735, South Plainfield, N.J. 07080
- Supplied by Mitsubishi Corporation UK Plc., Bow Bells House, Bread Street, London, EC4M 9BQ, UK
- 5. Supplied by US Cosmetics, 313 Lake Rd, Dayville, CT 06241, USA
- 6. Supplied by Chesham Chemicals Ltd., Cunningham House, Westfield Lane, Kenton, Harrow, Middlesex, UK
- Supplied by BASF, Earl Road, Cheadle Hulme, Cheadle, Cheshire, SK8 6QB

Method of Preparation

The above compositions can be prepared as follows:

The ingredients of phase A are mixed together for approximately five minutes with shear mixing until homogenised. With high speed shear mixing the materials of phase B are added gradually to phase A and then the batch is mixed for twenty minutes. Phase C is then added and the resulting mixture is ground until fully dispersed. Wax phase D is then added and the batch is heated to 85°C in a water bath. When the waxes have melted, the batch is removed from the water bath and cooled to 50°C. A pre-mix of phase E is then added to the batch and homogenised with a high shear mixer for 10 minutes. The batch is then cooled to room temperature. A pre-mix of phase F is added to the batch and homogenised for 10 minutes. Then phase G is added to the batch and homogenised for 10 minutes. The water phase is made by adding all ingredients together and mixing until all salts/powders have dissolved. The water phase is added to the oil phase quickly whilst homogenising at a low speed with stirring. When all of the water phase had been added, high shear is applied to the batch for approximately 5 minutes to increase the viscosity of the final product. The resulting make-up composition is ready for packaging.

The cosmetic compositions of the Examples exhibit significant reduction in the appearance of fine lines and wrinkles, together with improved coverage and skin feel and application characteristics.

What is claimed is:

- 1. Cosmetic composition comprising:
 - (a) from about 0.01% to about 20% by weight of particulate nylon; and
 - (b) from about 0.01% to about 20% by weight of a liquid, polyol carboxylic acid ester having a polyol moiety and at least 4 carboxylic acid moities, wherein the polyol moiety is selected from sugars and sugar alcohols containing from about 4 to about 8 hydroxyl groups, and wherein each carboxylic acid moiety has from about 8 to about 22 carbon atoms, and wherein said liquid polyol carboxylic acid ester has a complete melting point of less than about 30°C.
- 2. Cosmetic composition according to Claim 1 wherein the particulate nylon has a volume average particle size in the range of from about 5 microns to about 30 microns, preferably from about 10 microns to about 20 microns.
- Cosmetic composition according to Claim 1 or 2 wherein the nylon is nylon 12.
- 4. Cosmetic composition according to any of Claims 1 to 3 wherein said liquid polyol carboxylic acid ester contains no more than about 2 free hydroxyl groups.
- 5. Cosmetic composition according to any of Claims 1 to 4 wherein said carboxylic acid moities contain from about 14 to about 18 carbon atoms.
- 6. Cosmetic composition according to any of Claims 1 to 5 wherein said polyol moiety is selected from erythritol, xylitol, sorbitol, glucose, sucrose, and mixtures thereof.
- 7. Cosmetic composition according to any of Claims 1 to 6 wherein said polyol moisty is sucrose.

- 8. Cosmetic composition according to any of Claims 1 to 7 wherein said liquid polyol carboxylic acid ester has a complete melting point below about 27.5°C.
- 9. Cosmetic composition according to any of Claims 1 to 8 wherein said liquid polyol carboxylic acid polyester has a complete melting point below about 25°C.
- 10. Cosmetic composition according to any of Claims 1 to 9 wherein said liquid carboxylic acid polyol ester is selected from sucrose pentaoleate, sucrose hexaoleate, sucrose heptaoleate, sucrose octaoleate, and mixtures thereof.
- 11. Cosmetic composition according to any of Claims 1 to 10 comprising from about 4% to about 8% by weight of the particulate nylon.
- 12. Cosmetic composition according to any of Claims 1 to 11 comprising from about 1% to about 10% by weight of the liquid, polyol carboxylic acid ester.
- 13. Cosmetic composition according to any of Claims 1 to 12 wherein the composition is in the form of a water-in-oil emulsion comprising a continuous oil phase and a discontinuous aqueous phase.
- 14. Cosmetic composition according to Claim 13 wherein the oil phase comprises from about 0.01% to about 25% by weight of oil phase of non-volatile silicones and from about 75% to about 85% by weight of the oil phase of volatile silicones.
- 15. Use of a cosmetic composition according to any of Claims 1 to 14 reducing the appearance of wrinkles and/or fine lines.

INTERNATIONAL SEARCH REPORT

PCT/IB 98/00462

	SIFICATION OF SUBJECT MATTER			
IPC 6				
According	to International Patent Classification (IPC) or to both national or	tassification and IPC		
	S SEARCHED			
IPC 6	documentation searched (classification system followed by clas A61K C08K	stication symbols)		
Document	tation searched other than minimum occumantation to the extent	that such documents are included in the lierds s	earched	
Electronic	data base consulted during the international search (name of d	ala pase and, where practical, search terms use	d)	
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT			
Category *		he relevant passages	Relevant to claim No.	
A	EP 0 170 000 A (CHEESEBROUGH-I 5 February 1986 see claims 1,2; examples 2B,20	1		
A	DATABASE "CHEMICAL ABSTRACTS" Abs. 120: 116 515, Colombus, (& JP 05 279 236 A (SHISEIOO CO 26 October 1993 XP002070510	1		
A	DATABASE "CHEMICAL ABSTRACTS" Abs. 124: 66 240, Colombus, Option of the Colombus of the Colomb	1		
Furt	ther documents are listed in the continuation of box C.	Patent family members are listed	in annex.	
"A" docume consider a docume which catatio. "O" docume other in "P" docume later to Date of the	ent defining the general state of the art which is not pered to be of particular relevance documents but published on or after the international state and which may throw doubts on pnority claim(s) or is cited to establish the publicationdate of another in or other special reason (as specialed) ent referring to an oral disclosure, use, exhibition or means and provided to the international filling date but here the priority date claimed. July 1998	To later document published after the interpretation of priority date and not in conflict with cated to understand the principle or the invention. "X" document of particular relevance; the cannot be considered novel or cannot involve an inversive step when the document of particular relevance; the cannot be considered to involve an indocument is combined with one or more ments, such combination being obvious in the art. "&" document member of the same patent. Data of marting of the international sea.	mational filing data the application but acry underlying the daimed invertion the considered to cument to taken alone daimed invention ventive step when the ara other such docu- tes to a person skilled family	
Name and r	making address of the ISA European Parent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tal. (+31-70) 340-2040, Tx. 31 651 epo nt, Fax: (+31-70) 340-3016	Authorized efficer Glikman, J-F		

INTERNATIONAL SLARCH REPORT

Information of restort family members

PCT/IS 98/00462

Patent document cited in search report		Publication gate	Patent family member(s)		Publication date
EP 170000	Α	05-02-1986	US	4712571 A	15-12-1987
			AU	566043 8	08-10-1987
			AU	4427785 A	02-01-1986
			BR	8502553 A	04-02-1986
			CA	1269940 A,C	05-06-1990
			DK	149992 A	14-12-1992
			DK	296185 A	30-12-1985
			FI	851547 A	30-12-1985
			ΙE	58021 8	16-06-1993
			IN	165831 A	20-01-1990
			JP	1768845 C	30-06-1993
			JP	4045485 B	27-07-1992
			JP	61015814 A	23-01-1986
			CH	674459 A	15-06-1990

Form PCT/ISA/210 (patent family arries) (Auty 1992)